PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLIST	HED (UNDER THE PATENT COOPERATION TREATY (PCT)			
(51) International Patent Classification ⁵ :		(11) International Publication Number: WO 94/21668			
C07K 1/08, 5/08	A1	(43) International Publication Date: 29 September 1994 (29.09.94)			
(21) International Application Number: PCT/US (22) International Filing Date: 23 March 1994 (23) (30) Priority Data: 08/036,378 24 March 1993 (24:03:93) (71) Applicant: THE DU PONT MERCK PHARMACE COMPANY [US/US]; 1007 Market Street, Wilmin 19898 (US). (72) Inventor: KETINER, Charles, Adrian; 2411 Chatha Wilmington, DE 19803-2709 (US). (74) Agents: REINERT, Norbert, F. et al.; The Du Por Pharmaceutical Company, Legal/Patent Records 1007 Market Street, Wilmington, DE 19898 (US).	23.03.9 UTICA gton, D m Driv the Merces Centes	(81) Designated States: AU, BB, BG, BR, BY, CA, CN, CZ, FI, GE, HU, JP, KG, KP, KR, KZ, LK, LV, MD, MG, MN, MW, NO, NZ, PL, RO, RU, SD, SI, SK, TJ, TT, UA, UZ, VN, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG). Published With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.			
		·			
(54) Title: REMOVAL OF BORONIC ACID PROTECT	ING GI	ROUPS BY TRANSESTERIFICATION			
(57) Abstract		·			

A method for the removal of ester protecting groups from α -amino boronic acid is disclosed for the preparation of compounds of the formula: R^1 - X_{cr} NHCH(R^2)-B(OH)₂.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	GB	United Kingdom	MIR	Mauritania
ΑŪ	Australia	GE	Georgia	MW	Malawi
BB	Barbados	GN	Guinea	NE	Niger
BE	Belgium	GR	Greece	NL	Netherlands
BF	Burkina Faso	HU	Hungary	NO	Norway
BG	Bulgaria	IΕ	Ireland	· NZ	New Zealand
BJ	Benin	. 11	Italy	PL	Poland .
BR	Brazil	JP.	Japan	PT	
BY	Belarus	KE	Kcuya	RO	Portugal
CA	Canada	KG	Kyrgystan	RU .	Romania
CF	Central African Republic	KP	D		Russian Federation
CG	Congo		of Korea	- SD	Sodan
CH	Switzerland	KR	Republic of Korea	SÉ	Sweden
a	Côte d'Ivoire	KZ	Kazakhstan	SI	Slovenia
CM	Cameroon	Ľ	Liechtenstein	SK	Slovakia
CN	China	LK		SN	Scocgal
CS CS	Czechoslovakia	LU	Sri Lanka	TD	Chad
cz	Czech Republic		Luxembourg	10	Togo
DE	Gennary	.LV	Latvia	IJ	Tajikistan
DK	Denmark	MC	Monaco	II	Trinidad and Tobago
		MD	Republic of Moldova	UA	Ukraine
ES	Spain	MG	Madagascar	US	United States of America
Ħ	Finland	ML	Mali	UZ	Uzbekistan
FR	France	MN	Mongolia	VN	Vict Nam
GA	Gabon				

-1-

Title

Removal of Boronic Acid Protecting Groups by Transesterification

5

10

Field of the Invention

The present invention relates to a process for the removal of ester protecting groups from α -amino boronic acids and corresponding peptide analogs by transesterification with hydrophobic boronic acids.

Background of the Invention

Simple boronic acids are inhibitors of serine proteases. For example, Koehler et al. Biochemistry 10: 2477 (1971) reports that 2-phenylethane boronic acid 15 inhibits chymotrypsin at millimolar levels. synthesis of boronic acid analogs of N-acyl-a-amino acids has yielded more effective inhibitors. AcboroPhe-OH, R-1-acetamido-2-phenylethane boronic acid, inhibits chymotrypsin with a K_i of 4 μM Matteson et al. 20 J. Am. Chem. Soc. 103: 5241 (1981). More recently, Shenvi, US 4,537,773 (1985) disclosed that boronic acid analogs of α -amino acids, containing a free amino group, were effective inhibitors of aminopeptidases. Shenvi, US 4,499,082 (1985) discloses that peptides containing 25 an α -amino boronic acid with a neutral side chain were more effective inhibitors of serine proteases exceeding inhibitors disclosed earlier by as much as 3 orders of magnitude in potency. The chemistry of α -aminoboronic acids was further expanded to the synthesis of peptide. 30 analogs containing boronic acid with positive charged side chains, boroLysine, boroArginine, boroOrnithine, and isothiouronium analogs. This is disclosed in Kettner, et al. EPA 0,293,881, published December 7, 1988. 35

Much progress has been made in the synthesis of boronic acid and corresponding peptides with the

boronic acid protected as an ester. However, a convenient method of removal of the ester protecting group is lacking. Matteson (1981) infra, reports the destructive removal of pinanediol group by treatment with anhydrous BCl3. Kettner and Shenvi J. Biol. Chem. 15106 (1984) describe the removal of the pinacol protecting group by converting the boronic pinacol esters to the thermodynamically more stable, diethanolamine ester by transesterification and then 10 hydrolysis by treatment with aqueous acid or with a cation exchange resin. This method is not applicable for removal of pinanediol ester due to its greater stability. Matteson Chem. Rev. 89: 1535 (1989) describes the removal of the pinanediol group in situ by 15 incubations in borate buffer. It should be noted that the pinanediol ester is preferred in synthesis due to it ability to direct stereochemistry at the α -carbon of boronic acid and its stability to chemical manipulations. The pinanediol protecting group was used 20 almost exclusively in the preparation of boroArginine peptides, shown in EPA 0,293,881. In one example, partial hydrolysis of the pinanediol ester was obtained by binding Ac-(D)Phe-Pro-boroArg-C10H16 to a cation exchange resin and washing extensively with aqueous acetic acid followed by elution with HCl. This reaction is slow, it requires recovery of product by evaporation of large volumes of water and separation of the free boronic acid from the ester . Removal of the pinanedial by treatment with BCl3 as the final step in synthesis

35

Summary of the Invention

.. .

The present invention provides a method for converting compounds of formula I

30 was considered to be the only practical method.

:. ·.

.

and the second of the second of the second

PCT/US94/02964

$$R^1-X_n-NHCH(R^2)-BR^3R^4$$
(1)

5 to compounds of formula II,

$$R^1-X_n$$
-NHCH (R^2) -B (OH) ₂
(II)

(11)

wherein for both formula I and formula II $10 \quad R^1$ is

- a) hydrogen,
 - b) an N-terminal protecting group,
- c) $-SO_2(CH_2)_m$ -aryl, wherein aryl is phenyl, napthyl or biphenyl substituted with one, two or three
- substituents selected from the group consisting of halo (F, Cl, Br, I,), -CN, Cl-Cl0-alkyl, C3-C8-cycloalkyl, C2-Cl0-alkenyl, C2-Cl0-alkynyl, -OR 7 , -NO $_2$, -CF $_3$, -S(O) $_r$ R 8 , -NR 6 R 7 , -COR 7 , -CO $_2$ R 7 , -CONR 6 R 7 ;

X is a peptide of 1-20 amino acids;

- $20 R^2 is$
 - a) C1-C10-alkyl,
 - b) C2-C10-alkyl-Y,
 - c) -(CH₂)_n-aryl, wherein aryl is as defined above;

Y is

- a) $-NHC(NH)NH_2$,
 - b) $-NH_2$,
 - c) $-SC(NH)NH_2$,
 - d) $-OR^9$,
 - e) -SR⁹;
- 30 R^3 and R^4 are
 - a) C1-C8-alkoxy, or
 - b) when taken together \mathbb{R}^3 and \mathbb{R}^4 form a cyclic boronic ester where said chain or ring contains from 2 to 20 carbon atoms and, optionally, a heteroatom which
- 35 can be N, S, or O; R⁵ and R⁶ are independently
 - a) H,

r is 0 to 2;

```
b) C1-C8-alkyl,
              c) C1-C8-alkoxy,
              d) C3-C8-cycloalkyl,
              e) -CO_2R^7
  5
              f) - (CH<sub>2</sub>)<sub>m</sub>-phenyl;
       R^7 is
             a) H,
             b) phenyl,
             c) benzyl,
 10
             d) C1-C8-alkyl;
      R<sup>8</sup> is
             a) phenyl,
             b) C1-C4-alkyl,
             c) C1-C4-alkoxy,
15
             d) -CF3;
      \mathbb{R}^9 is
             a) H,
             b) C1-C2-alkyl,
             c) phenyl or phenyl optionally substituted with a
      substituent selected from the group consisting of halo
20
      (F, Cl, Br, I), -CN, Cl-Cl0-alkyl, C3-C8-cycloalkyl, C2-
     C10-alkenyl, -C2-C10-alkynyl, -OR^7, -NO_2, -CF_3,-S(O)_rR^8,
     -NR<sup>6</sup>R<sup>7</sup>, -COR<sup>7</sup>, -CO<sub>2</sub>R<sup>7</sup>, -CONR<sup>6</sup>R; ^7 wherein R<sup>5</sup>, R<sup>6</sup>, and R<sup>8</sup>
     are as defined above;
     n is 0 or 1;
25
     m is 0 to 2;
```

which comprises reacting a compound represented by

formula I in a mixture of water and a water-immiscible
organic solvent containing an organic boronic acid
acceptor present in an amount equal to at least 1
equivalent of the compound of formula I, stirring the
mixture at a temperature in a range of from about 5 to

about 35°C, for a time of approximately 1 hour, allowing
the mixture to then separate into two distinct phases,

separating the phases and then recovering the desired compound of formula II from the separated aqueous phase.

5 <u>Detailed Description of the Invention</u>

As used throughout the specifications, the following abbreviations for amino acid residues or amino acids apply:

Ala = L-alanine

10 Arg = L-arginine

Asn = L-asparagine

Asp = L-aspartic acid

Cys = L-cysteine

Gln = L-glutamine

15 Glu = L-glutamic acid

Gly = glycine

His = L-histidine

Ile = L-isoleucine

Leu = L-leucine

20 Lys = L-lysine

Met = L-methionine

Phe = L-phenlyalanine

Pro = L-proline

Ser = L-serine

25 Thr = L-threonine

Trp = L-tryptophan

Tyr = L-tyrosine

Val = L-valine

The "D" prefix for the foregoing abbreviations indicates
30 the amino acid is in the D-configuration. "D,L"
indicates the amino is present in mixture of the D- and
the L-configurations.

Other abbreviations used throughout the description

35 below

are:

Me

= methyl

35

```
Εt
                     = ethvl
          Boc
                     = t-butoxycarbonyl
          Z
                     = benzyloxycarbonyl
          2Clz
                     = 2-chlorobenzyloxycarbonyl
 5
          4Clz
                     = 4-chlorobenzyloxycarbonyl
          p-N0_2-Z
                     = p-N0<sub>2</sub>benzyloxycarbonyl
          AC ·
                     = acetyl
                     = adamantyloxycarbonyl
          Adc
          DIPA
                     = diisopropylamine
10
          DIPEA
                     = diisopropylethylamine
          DCHA
                     = dicyclohexylamine
          DBU
                     = 1,8-diazabicyclo[5.4.0]undec-7-ene
          DABCO
                     = 1,4-diazabicyclo[2.2.2]octane
          MMM
                     = N-methylmorpholine
15
          DMAP
                     = 4-dimethylaminopyridine
          FSA
                     = formamidinesulfinic acid
          FAB/MS
                     = fast atom bombardment mass
    spectrometry
                                     MS(NH_3-C1) = chemical
    ionization mass spectrometry
20
          NMR
                     = nuclear magnetic resonance
    spectrometry
```

The following reagents were obtained from commercial sources: l-hydroxybenzotriazole•H₂O, adamantylfluoroformate, di-t-butyldicarbonate, benzyloxycarbonyl chloride, 2-chlorobenzyloxycarbonyl chloride, N-hydroxysuccinimide, formamidinesulfinic acid, 32% peracetic acid.

Boc-Pro-boroOrn-C₁₀H₁₆, Ac-(D)Phe-Pro-boroOrn-C₁₀H₁₆, BocPhe-boroOrn-C₁₀H₁₆ benzenesulfonic acid were prepared by the procedure described in EP0293881A2, pl2-13.

The prefix "boro" indicates amino acid residues where the carboxy group is replaced by a boronic acid (formula II, \mathbb{R}^3 and \mathbb{R}^4 = -OH).

The pinanediol boronic acid ester and the pinacol boronic acid ester are abbreviated "- $C_{10}H_{16}$ " and " $C_{6}H_{12}$ ",

WO 94/21668 PCT/US94/02964

> respectively. Other illustrations of diols useful for deriving a boronic acid esters are 1,2-ethanediol, 1,3propanediol, 2,3-butanediol, 1,2-diisopropylethanediol, 5,6-decanediol, 1,2-dicyclohexylethanediol.

-7-

Note that throughout the text when an alkyl substitutent is mentioned, the normal alkyl structure is meant (e.g. butyl is n-butyl) unless otherwise specified. However, in the definition of radicals above (e.g. R²), both branched and straight chains are included in the scope of alkyl.

10

15

It is understood that many of the compounds of the present

invention contain one or more chiral centers and that these stereoisomers may possess distinct physical and biological properties. The present invention comprises all of the stereoisomers or mixtures thereof. If the pure enantiomers or diastereomers are desired, they may be prepared using starting

15 materials with the appropriate stereochemistry, or may be separated from mixtures of undesired 20 stereoisomers by standard techniques, including chiral chromatography and recrystallization of diastereomeric salts.

"N-terminal protecting group" as used herein, 25 refers to various art recognized amino-terminal protecting groups employed in peptide synthesis. Examples of suitable groups include formyl, acetyl, benzoyl, trifluoroacetyl, and methoxysuccinyl; aromatic urethane protecting groups, such as, benzyloxycarbonyl; and aliphatic 30 urethane protecting groups, such as t-benzyloxycarbonyl or adamantyloxycarbonyl. Gross and Meinhoffer, eds., The Peptides, Vol. 3; 3-88 (1981), Academic

Press, New York 1981, disclose numerous suitable amine protecting groups and is incorporated herein by reference for

that purpose.

"Peptide of 1-20 amino acids" as used herein, peptide chain of one to twenty natural or unnatural 5 amino acids of either D- or L-configuration. Roberts and Vellaccio, The Peptides, Vol. 5; 341-449, Academic Press, New York 1983, disclose numerous suitable natural and unatural amino acids and is incorporated herein by reference for that purpose. This term is also intended 10 to include sidechain protected amino acid residues that are commonly employed in peptide synthesis such as those disclosed in the Peptides, Vol 3, 3-88 (1981). This reference is incorporated herein by reference for that 15 purpose.

It should be noted that to yield a compound of formula II where X is a peptide, optionally, the N-terminal or sidechain protecting groups can be removed by using procedures well known to those skilled in the art. For example, where the N-terminal or side chain protecting group is BOC, the BOC group can be removed by treatment with Anhydrous HCL. Where the N-terminal or side chain protecting group is Z, the Z group can be removed by means of catalytic hydrogenation.

25

30

20

The present invention relates to the synthesis of free boronic acids (compounds of formula II) from ester precursors by transesterification reactions with aliphatic and aromatic boronic acids under heterogeneous reaction conditions.

Mark Congressing. Geography

Scheme 1

10

15

20

This novel method is readily applicable to compounds where the R² side chain is positively charged as shown in Scheme 1 where R² is the 3-quanidino-propyl moiety. In this example, the protected boronic acid ester, Ac-(D) Phe-Pro-boroArg-C10H16, is suspended in a mixture consisting of water, an equal volume of diethyl ether, and 5 equivalents of phenyl boronic acid. The flask is stoppered and allowed to stir rapidly with a magnetic stirrer at room temperature. Two clear phases are observed after 15-30 min. Stirring is continued for 3 The reaction mixture is transferred to a separatory funnel where the phases are separated. The aqueous phase is then washed with two portions of ether. is removed by evaporation at 35-43°C at a reduced 25 pressure. Products are usually obtained as white foams after drying in vacuo, with KOH and P2O5 and are readily converted to amorphous white solids by triturating with ether.

10

boroArg-C₁₀ H₁₆

isothiouronium Salt Analog of boroArg-C₁₀ H₁₆

boroLys-C10 H16

H-boroVal-C₆H₁₂

MeOSuc-Ala-Ala-Pro-(D,L)boroVal-C6H12

The above process depends on the final product being more soluble in the aqueous phase than the organic phase. This criteria is readily met for compounds such as the boroArginine, boroLysine, and boroOrnithine peptides as well as analogs were the isothiouronium group replaces the guanidino group. It is applicable to compounds in US 4,537,773 and US 4,499,082 which describe α-aminoboronic acids with neutral side chains

WO 94/21668 PCT/US94/02964

-11-

and peptides containing α -aminoboronic acids with neutral side chains, respectively. For removal of the ester protecting group from α -aminoboronic such as HboroVal-C6H12, this method should be generally applicable since these compounds are readily soluble in water due to the presence of the free α -amino group. should be applicable to a large number of less hydrophobic peptide boronic acids which are readily soluble in water. For example, the pinacol protecting group of MeOSuc-Ala-Ala-Pro-boroVal-OH is readily 10 removed by the method of the present invention. However, it will be desirable to run trial reactions on a small scale to determine the solubility of the free boronic acid product and the feasibility of this method. For more hydrophobic compounds in this series, it maybe 15 necessary to design a synthetic protocol were the transesterification step is applied to intermediates containing charged residues.

The use of a biphasic system with the organic phase consisting of diethyl ether and phenyl boronic appears to be ideal for the preparation of most free boronic acids. This method will be applicable to the removal of other boronic acid protecting groups represented by R³ and R⁴ in formula (I). Specific examples, in addition to the pinanediol and pinacol groups, are where R³ and R⁴ taken together form a moiety derived from 1,2-ethanediol, 1,3-propanediol, 2,3-butanediol, 1,2-diisopropylethanediol, 5,6-decanediol, or 1,2-dicyclohexylethanediol. The protecting groups can also be where R³ and R⁴ are derived from alcohols such as isopropanol, methanol, ethanol or n-propanol. Of course, R³ and R⁴ can each be derived from the same alcohol or from different alcohols, if desired.

20

30

Organic solvents other than diethyl ether can be used in the method of the invention. It is only necessary that the organic solvent be water immiscible. Suitable choices of other organic solvents are

5

10

carbonteterachloride, chloroform, methylenechloride, ethyl acetate, benzene, tolulene or hexane.

Boronic acid acceptors for the ester protecting group other than phenyl boronic acid also can be used in the method of the invention. It is only necessary that the acceptor boronic acid, both in its free form and in its esterified form, have greater solubility in the organic phase than in the aqueous phase. Suitable choices of other acceptor boronic acids are butyl boronic acid, pentyl boronic acid, hexyl boronic acid or cyclohexyl boronic acid.

For the method of invention, the ratio of water to organic solvent in the mixture in which the ester precursor of formula (I) is suspended can vary widely.

- It is important that sufficient volumes of water and organic solvent be present to completely dissolve the products of the reaction (acceptor boronic acid plus ester for the organic phase and free boronic acid for the aqueous phase).
- For the method of the invention, the amount of acceptor boronic acid in the reaction mixture should be an amount equal to at least a molar equivalent of the ester precursor of formula (I) present in said mixture. Generally, it is preferable to have the acceptor boronic
- acid present in an amount in excess of an equimolar
 amount, the most preferred amount-being a range of from 3 to 5 equivalents.

The time of stirring the reaction mixture can vary over wide limits depending on the ester precursor and the acceptor boronic acid involved. Usually, the minimum time for stirring is 1 hour, but can vary from 0.2 to 48 hours.

The second of th

In the method of the invention, the desired

35 product compound of formula (II) is recovered from the aqueous phase after its separation from the two phase system formed from stirring the reaction mixture. This

A CONTRACTOR OF A STATE OF THE STATE OF THE

WO 94/21668 PCT/US94/02964

-13-

is best accomplished by the removal of water from the aqueous phase by means well understood by those skilled in the art, such as with a rotary evaporator.

NMR, proton nuclear magnetic resonance, chemical shifts are reported in δ units, parts per million 5 downfield from the internal tetramethylsilane standard. Elemental analyses were conducted by Galbraith Laboratories Inc., Knoxville, TN and Microanalysis Inc., Wilmington, DE. FAB/MS samples of free boronic acids did 10 not give consistent results making it difficult to monitor the removal of ester protecting groups difficult by this means. However, the presence of the pinanediol and the pinacol groups are readily observed in NMR spectra. For the pinanediol ester, a methyl group is observed at $\partial 0.9$ and the methyl groups of the pinacol 15 groups are observed as singlet at δ 1.1 Following the removal of pinanediol protecting group, FAB/MS were run by treating the sample with ~2 equivalents of pinacol in methanol for 5 min and evaporating the solvent.

20 Similarly, FAB/MS samples of free boronic acid, obtained by removal of the pinacol, were prepared by treating with pinanediol.

Example 1

Preparation of Ac-(D)Phe-Pro-boroArg-OH*benzene,
 sulfonic acid.

30

35

The synthesis of Ac-(D)Phe-Pro-boroArg-C₁₀H₁₆•benzene sulfonic acid has been described previously, Kettner et al. *J. Biol Chem* **265**: 18289 (1990).

Ac-(D)Phe-Pro-boroArg-C₁₀H₁₆•benzene sulfonic acid (0.20 g, 0.27 mmoles) and phenyl boronic acid (0.16 g, 1.3 mmoles) were suspended in a mixture consisting of 5 ml of water and 5 ml of ether. The mixture was stirred overnight at room temperature. The two phases were separated, the organic phase was washed with water, and the aqueous phase was washed with ether. The combined

aqueous phases was evaporated to vield 0.14 g of product. NMR was consistent with the desired structure and the product obtained in Example 2.

5

Example 2

Preparation of Ac-(D)Phe-Pro-boroArg-OH·HCl, Ac-(D)Phe-Pro-boroArg- $C_{10}H_{16}$ •benzene sulfonic acid (6.4 g, 8.5 mmoles) and phenyl boronic acid (5.2 g, 42)mmoles) were suspended in 150 ml of water and 150 ml of ether. The mixture was stirred overnight. The phases 10 were separated and the ether phase was washed with two 100 ml portions of water. The combined aqueous phases were washed with ether. The aqueous phase was concentrated to ~ 50 ml by evaporation and then it was passed through a column containing 15 ml of BioRadTM 15 AG1-X8 (Cl form). The aqueous phase was further concentrated to ~2 ml and it was chromatogramed on a 2.5 x 100 cm column containing BioRad TM P2 resin and equilbrated with 1.0 mM HCl. Fractions containing the desired product were pooled, evaporated, dried in vacuo. 20 and triturated with ether to yield 3.4 g. Anal. Calcd. for $C_{21}H_{34}N_{6}O_{5}BC1$: C=50.77%,

H=6.91%, N=16.92%, and B=2.18%. Found: C=50.91%, H=6.97%, N=16.91%, B=2.29%.

25

Example 3

Preparation of Ac-Phe-Pro-boroArg-OH•HCl 30 The starting material for this reaction, Ac-Phe-Pro-boroArg-C10H16 • HCl, was prepared by coupling Ac-Phe-OH to H-Pro-boroArg-C10H16. The intermediate Boc-ProboroOrn- $C_{10}H_{16}$ was prepared by the procedure described in EPA 0 293 881 and it was guanidated using 35 aminoiminomethane sulfonic acid [Mosher et al.

Tetrahedral Lett. 29: 3183 (1988)]. Boc-Pro-boroOrn-C10H16 benzene sulfonic acid (4.8 g, 10.4 mmoles) was

dissolved in 50 ml of absolute ethanol; 4-dimethylaminopyridine (2.5 g, 20.7 mmoles) and aminoiminomethane sulfonic acid (2.6 g, 20.7 mmoles) were added. The mixture was refluxed at 80°C for 3 hrs

were added. The mixture was refluxed at 80°C for 3 hrs.

It was cooled and solids were removed by filtration.

Solvent was evaporated, the residue was dissolved in chloroform, and it was washed with 0.2 N HCl prepared in saturated aqueous NaCl and with saturated aqueous NaCl. After drying over anhydrous sodium sulfate, solvent was evaporated to yield 5.4 g of a foam. This material was dissolved in methanol and it was chromatogramed on a 2.5 x 100 cm column of SephadexTM LH-20 using methanol as a solvent. Product, 4.4 g, was obtained. FAB/MS calcd. for M (C25H44N5O5B) + H: 506.56. Found: 506.49.

H-Pro-boroArg-C10H16•2HCl was prepared by dissolving Boc-Pro-boroArg-C10H16•HCl (1.3 g, 2.4 mmoles) in 10 ml of dioxane and adding 10 ml of 3.3 N HCl: dioxane. After stirring for 2 hrs, solvent was evaporated and the residue was triturated with ether to yield 1.2 g of product. FAB/MS calcd. for M (C20H36N5O3B) + H: 406.43. Found: 406.38.

Ac-Phe-OH (87 mg, 0.42 mmoles) was coupled to H-Pro-boroArg-C10H16•2HCl (200 mg, 0.42 mmoles) using the carbodiimide procedure. The starting materials were dissolved in 20 ml of methylene chloride, N-methylmorpholine (92 μl, 0.84 mmoles), 1-hydroxybenzotriazole•H₂O (130 mg, 0.84 mmoles), and dicyclohexylcarbodiimide (86 mg, 0.42 mmoles) were added. After stirring overnight at room temperature, the reaction mixture was filtered, the filtrate evaporated, and the residue was chromatogramed 2.5 x 50 cm column of LH-20 using methanol as a solvent. The desired product was obtained in a yield of 240 mg.

Ac-Phe-Pro-boroArg- $C_{10}H_{16} \cdot HC1$ (0.13 g, 0.21 mmoles) and phenyl boronic acid (0.13 g, 1.0 mmoles)

35

595.41.

FAB/MS calcd. for M (C31H47N6O5B) + H: 595.66. Found:

were dissolved in a mixture of 5 ml of water and 5 ml of ether. The mixture was stirred 3 hrs at room temperature. The reaction phases were separated and the aqueous phase was extensively washed with ether. Water was evaporated and the residue dried to yield 0.11 g. The product was triturated with ether to yield a white solid. FAB/MS calcd. for the pinacol ester, M $(C_{27}H_{43}N_{6}O_{5}B)$ + H: 543.58. Found: 543.48.

10

Example 4

Preparation of Ac-Pro-boroArg-OH•HCl
Ac-Pro-boroArg-C10H16•HCl was prepared by
dissolving H-Pro-boroArg-C10H16•2HCl (200 mg, 0.41
mmoles) in 1 ml of dioxane: water (1:1) and adding
acetic anhydride (59 µl, 0.63 mmoles) and sodium
bicarbonate (110 mg, 1.2 mmoles). The reaction was
allowed to stir 30 min at room temperature, it was
acidified with HCl, diluted with methanol, and
evaporated. It was redissolved in methanol and
chromatogramed on 2.5 x 50 cm column of LH-20.
Fractions containing the desired product were pooled,
evaporated, and triturated with ether to yield 140 mg.
FAB/MS calcd. for M (C22H38N5O4B) + H: 447.97. Found:
448.43.

The conditions in Example 3 were used to prepare the free boronic acid of Ac-Pro-boroArg-C10H16·HCl (0.12 g, 0.24 mmoles). After triturating the product with ether, 0.080 g of Ac-Pro-boroArg-OH·HClwere obtained. FAB/MS calcd. for the pinacol ester, M (C18H34N5O4B) + 396.39. Found: 396.3.

Example 5

Preparation of Ac-Gly-boroArg-OH•benzene sulfonic acid
Boc-Gly-boroArg-C10H16 (10.2 g) was prepared from
Boc-Gly-boroOrn-C10H16•benzene sulfonic acid (12.5 g,
21.5 mmoles) by the procedure described in EPA 0 293

er de la companya de

.

30

881. FAB/MS calcd. for M $(C_{22}H_{40}N_{5}O_{5}B) + H$: 466.32 Found: 466.59.

 $H-Gly-boroArg-C_{10}H_{16} \cdot HCl$, benzene sulfonic acid was prepared by deblocking Boc-Gly-boroArg-C10H16 with HCl: dioxane.

Ac-Gly-boroArg-C₁₀H₁₆•benzene sulfonic acid was prepared by the procedure described for Ac-Pro-boroArg-C₁₀H₁₆ in Example 4. FAB/MS calcd. for M (C₁₉H₃4N₅O₄B) + H: 407.90. Found: 408.36.

The condition described for Example 3 were used to prepare the free boronic acid. Ac-Gly-boroArg-C10H16*benzene sulfonic acid (0.064 g, 0.11 mmoles) yielded 33 mg of Ac-Gly-boroArg-OH*benzene sulfonic acid. FAB/MS calcd. for the pinacol ester, M

(C15H30N5O4B) + H: 356.32. Found: 356.3.

Example 6

Preparation of Ac-(D)Phe-Gly-boroArg-OH•benzene sulfonic acid

Ac-(D)Phe-Gly-boroArg-C10H16 benzene sulfonic acid was prepared by coupling Ac-(D)Phe-OH to H-Gly-boroArg-C10H16 using a modification of the carbodimide procedure described in Example 3. For this coupling, 2 ml of dimethylformamide was used with 20 ml of methylene chloride as a solvent. FAB/MS calcd. for M (C28H43N6O5B) + H: 555.59. Found: 555.38.

The procedure described in Example 3 was used to prepare the free boronic acid. Ac-(D)Phe-Gly-boroArg-C10H16*benzene sulfonic acid (0.10 g, 0.14 mmoles) yielded 72 mg of Ac-(D)Phe-Gly-boroArg-OH*benzene sulfonic acid. FAB/MS calcd. for the pinacol ester M (C24H39N6O5B) + H: 503.51. Found: 503.32.

Example 7

Preparation of Boc-(D)Phe-Gly-boroArg-OH•HCl
Boc-(D)Phe-Gly-boroArg-C₁₀H₁₆ was prepared by
coupling Boc-(D)Phe-OH to the dipeptide analog using the

The control of the co

mixed anhydride procedure. The mixed anhydride of Boc-(D)Phe-OH (95 mg, 0.36 mmoles) was prepared by dissolving the acid in 3 ml of anhydrous tetrahydrofuran and adding N-methylmorpholine (40 μ l, 0.36 mmoles), and isobutyl chloroformate (46 μ l, 0.36 mmoles) at -20 $^{\circ}$ C. After 5 min, triethylamine (50 μ l, 0.36 mmoles) and 10 ml of cold tetrahydofuran were added and the mixture was immediately added to a 0°C solution of H-Gly-boroArg-C10H16•benzene sulfonic acid, HCl (200 mg, 0.36 mmoles) in 6 ml of chloroform. After allowing the reaction to 10 warm to room temperature and to stir several hrs, it was filtered and solvent was evaporated. The residue was chromatogramed on a 2.5 \times 50 cm column of LH-20 in methanol to yield 210 mg of the desired product. FAB/MS calcd. for M $(C_{31}H_{49}N_{6}O_{6}B) + H$: 613.39. Found: 15 613.65.

The procedure described in Example 3 was used to convert Boc-(D)Phe-Gly-boroArg-C10H16·HCl (0.050 g, 0.077 mmoles) to 36 mg of Boc-(D)Phe-Gly-boroArg-OH·HCl. FAB/MS calcd. for the pinacol ester, M (C27H45N6O6B) + H: 561.60. Found: 561.4.

25

20

Example 8

Preparation of Ac-Phe-Gly-boroArg-OH•benzene sulfonic acid

Ac-Phe-Gly-boroArg-C10H16•benzene sulfonic acid was prepared by coupling Ac-Phe-OH to H-Gly-boroArg-C10H16 using the carbodiimide procedure described in Example 3. FAB/MS calcd for M (C24H39N6O5B). 503.51. Found: 503.3.

Ac-Phe-Gly-boroArg-C10H16*benzene sulfonic acid (0.075 g, 0.10 mmoles) was treated with phenyl boronic acid by the procedure in Example 3 to yield Ac-Phe-Gly-boroArg-OH*benzene sulfonic acid. FAB/MS calcd. for the

WO 94/21668 PCT/US94/02964

-19-

pinacol ester, M (C27H43N4O5B) + H: 515.48. Found: 515.3.

Example 9

5 Preparation of Ac-(D)Phe-Pro-boroLys-OH•benzene sulfonic acid

The intermediate, NH2-CH[(CH2)4Br]BO2C10H16*HCl was prepared by the procedure described for the analogous compound, NH2-CH[(CH2)3Br]BO2C10H16*HCl, in

EPA 0 293 88Q. Also by analogous reactions, Ac-(D)Phe-Pro-NH-CH[(CH2)4Br]BO2C10H16, Ac-(D)Phe-Pro-NH-CH[(CH2)4N3]BO2C10H16, and Ac-(D)Phe-Pro-NH-CH[(CH2)4NH2]BO2C10H16*benzene sulfonic acid (Ac-(D)Phe-Pro-boroLys-C10H16*benzene sulfonic acid) were prepared.

Ac-(D)Phe-Pro-boroLys-C10H16*benzene sulfonic acid (0.50 g, 0.76 mmoles) was treated with phenyl boronic acid by the procedure described in Example 3 to yield Ac-(D)Phe-Pro-boroLys-OH*benzene sulfonic acid (0.35 g). FAB/MS calcd. for the pinacol ester, M (C27H43N4O5B) +

20 H: 515.48. Found: 515.3.

30

Example 10

Preparation of the Isothiouronium Analog of Ac-(D)Phe-25 Pro-boroArg-OH

Ac-(D)Phe-Pro-NH-CH[(CH₂)3-S-C(NH)-NH₂]BO₂-C₁₀H₁₆·HBr. was prepared by the procedure described in EPA 0 293 881. The corresponding bromide was treated with thiourea to yield the desired produce as an amorphous white solid. Anal. Calcd. for C₃₁H₄7N₅SBBr: C=53.75%, H=6.85%, N=10.11%, B=1.56%. Found: C=53.18%, H=6.68%, N=9.47%, and B=1.50%. FAB/MS calcd. for the pinacol ester, M (C₃₁H₄6N₅SB) + H: 612.71. Found: 612.36.

Ac-(D)Phe-Pro-NH-CH[(CH₂)₃-S-C(NH)-NH₂]BO₂-C₁₀H₁₆·HBr (1.0 g, 1.4 mmoles) was allowed to react with phenyl boronic acid by the procedure in Example 3 to

yield 0.66 g of the desired product, Ac-(D)Phe-Pro-NH-CH[(CH₂)₃-S-C(NH)-NH₂]B(OH)₂•HBr. Anal. Calcd. for C₂1H₃3N₅O₅SBBr: C=45.17%, H=5.97%, N=12.55%, and B=1.93%. Found: C=44.78%, H=5.58%, N=12.23%, and B=1.85%. FAB/MS calcd. for the pinacol ester, M (C₂7H₄2N₅O₅BS) + H: 560.31. Found: 560.41.

Example 11

Preparation of MeOSuc-Ala-Ala-Pro-(D,L)boroVal-OH

The synthesis of MeOSuc-Ala-Ala-Pro-(D,L)boroVal
C6H12 has been described previously, Kettner and Shenvi

J. Biol. Chem. 259: 15106 (1984). The pinacol ester

(100 mg, 0.17 mmoles) was allowed to react with 5

equivalent of phenyl boronic acid using the conditions

described in Example 3. The aqueous phase was

evaporated to yield 92 mg of MeOSuc-Ala-Ala-Pro
(D,L)boroVal-OH. NMR indicated only a trace (<10%) of
the pinacol group remained. FAB/MS calcd. for the

pinanediol ester, M (C30H49N4O8B) + H: 605.65. Found:

605.4.

Example 12

Preparation of H-(D,L)boroVal-OH
H-(D,L)boroVal-C6H12*trifluoroacetic acid (100 mg,
0.32 mmoles), described in Kettner and Shenvi (1984) was
allowed to react with phenyl boronic acid by the
procedure in Example 3. H-(D,L)boroValOH*trifluoroacetic acid was obtained in a yield of 76
mg. NMR was consistent with the desired structure
indicating the complete absence of the pinacol group.
FAB/MS calcd. for the pinanediol ester, M (C14H26NO2B) +
H: 252.22. Found: 252.2.

Example 13

Preparation of hydrocinnamoyl-Pro-boroLys-OH benzene sulfonic acid.

Hydrocinnamoyl-Pro-boroLys-C10H16 benzene sulfonic acid was prepared by the general procedure described in

WO 94/21668 PCT/US94/02964

-21-

EPA 0 293 881 and was allowed to react with phenyl boronic acid by the procedure in Example 3. The desired product was obtained in a yield of 92%. MS calcd. for M(C19H30N3O4B)+H-2H2O: 340.0. Found: 340. Anal Calcd. for C35H50N3O7SB: c=62.96%, H=7.55%, N=6.29%, B=1.62%. Found: C=62.75%, H=7.47%, N=6.28%, B=1.64%.

What is Claimed is:

A method for the preparation of a compound of formula (II)

 R^1-X_n -NHCH (R^2) -B $(OH)_2$ (II)

wherein

- $10 R^1 is$
 - a) hydrogen,
 - b) an N-terminal protecting group,
 - c) $-SO_2(CH_2)_m$ -aryl, wherein aryl is phenyl, napthyl or biphenyl substituted with one, two or three
- substituents selected from the group consisting of halo (F, C1, Br, I,), -CN, C1-C10-alkyl, C3-C8-cycloalkyl, C2-C10-alkenyl, C2-C10-alkynyl, -OR 7 , -NO $_2$, -CF $_3$, -S(O) $_r$ R 8 , -NR 6 R 7 , -COR 7 , -CO $_2$ R 7 , -CONR 6 R 7 ;

X is a peptide of 1-20 amino acids;

- $20 R^2 is$
 - a) C1-C10-alkyl,
 - b) C2-C10-alkyl-Y,
 - c) -(CH2) n-aryl, wherein aryl is as defined above;

Y is

- 25 a) $-NHC(NH)NH_2$,
 - b) $-NH_2$,
 - c) $-SC(NH)NH_2$,
 - $d) OR^9$
 - e) -SR⁹;
- 30 R⁵ and R⁶ are independently
 - a) H,
 - b) C1-C8-alkyl,
 - c) C1-C8-alkoxy,
 - d) C3-C8-cycloalkyl,
- 35 e) $-CO_2R^7$,
 - f) (CH₂)_m-phenyl;

40 R⁷ is

WO 94/21668 PCT/US94/02964

-23-

- a) H,
- b) phenyl,
- c) benzyl,
- d) C1-C8-alkyl;

5 R⁸ is

- a) phenyl,
- b) C1-C4-alkyl,
- c) C1-C4-alkoxy,
- d) -CF3;

 $10 R^9 is$

- a) H,
- b) C1-C2-alkyl,
- c) phenyl or phenyl optionally substituted with a substituent selected from the group consisting of halo
- 15 (F, Cl, Br, I), -CN, Cl-Cl0-alkyl, C3-C8-cycloalkyl, C2-Cl0-alkenyl, C2-Cl0-alkynl, $-OR^7$, $-NO_2$, -CF3, $-S(0)_RR^8$, $-NR^6R^7$, $-COR^7$, $-CO_2R^7$, $-CONR^6R^7$;

n is 0 or 1;

20 m is 0 to 2;

r is 0 to 2;

comprising suspending a compound of the formula

25 $R^1-X_n-NHCH(R^2)-BR^3R^4$

·(I)

wherein R^1 , R^2 , X, Y, R^5 , R^6 , R^7 , R^8 , R^9 , n, m and r are as defined above; and

30 R^3 and R^4 are

- a) C1-C8-alkoxy, or
- b) when taken together R^3 and R^4 form a cyclic boronic ester where said chain or ring contains from 2 to 20 carbon atoms and, optionally, a heteroatom which can be N, S, or O; R^3 and R^4 , independently, are optionally, a heteratom which can be N, S, or O,

in a mixture of water and a water-immiscible organic solvent containing an organic boronic acid acceptor present in an amount equal to at least 1 equivalent of said compound of formula (I), stirring said system for approximately one hour before allowing the reaction mixture to separate into two distinct phases, separating the phases, and recovering the compound of formula (II) from

10

15

5

2. The method of claim 1 wherein the water-immiscible organic solvent is selected from the group consisting of diethyl ether, carbonteterachloride, chloroform, methylene chloride, ethyl acetate, benzene, toluene or hexane.

the separated aqueous phase.

- 3. The method of claim 2 wherein the organic boronic acid acceptor is phenyl boronic acid.
- 20 4. The method of anyone of claims 1 to 3 wherein the amount of organic boronic acid receptor present in the suspending step is in the range of 3 to 5 molar equivalents of the amount of the compound of formula (I) present in said step.

13.

25

5. The method of claim 1 wherein the compound of formula (II) is recovered from the seperated aqueous phase by the evaporation of water from said phase.

30

6. The method of claim 5 wherein the evaporation of water is by means of a rotary evaporator.

INTERNATIONAL SEARCH REPORT

Inter onal Application No

	<u> </u>	PC	CT/US 94/02964	
A. CLAS	SIFICATION OF SUBJECT MATTER CO7K1/08 CO7K5/08			
l .	to International Patent Classification (IPC) or to both national	dassification and IPC		
	OS SEARCHED documentation searched (classification system followed by classification system followed by cla	rsification symbols)	,	
IPC 5	СО7К	mucason vymmony		
Document	ation searched other than minimum documentation to the exten	t that such documents are included	in the fields searched	
Electronic	data base consulted during the international search (name of da	ta base and, where practical, search	h terms used)	
• •				
C. DOCUM	MENTS CONSIDERED TO BE RELEVANT			
Category *	Citation of document, with indication, where appropriate, of	the relevant passages	Relevant to claim No	٥.
A	EP,A,O 293 881 (E.I. DUPONT NE COMPANY) 7 December 1988	MOURS AND		
A	US,A,5 196 948 (D.H.KINDER ET April 1992	AL.) 21		
A .	CHEMICAL REVIEWS vol. 89 , 1989 , USA pages 1535 - 1551 D.S.MATTESON 'Alpha- Halo Boronic Esters: Intermediates for Stereodirected			
	Synthesis¹			
Furt	her documents are listed in the continuation of box C.	X Patent family member	ers are listed in annex.	
* Special cat	legories of cited documents :	T later document published	after the international liting date	
A document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on or after the international filing date "X" document of particular relevance; "E" carlier document but published on or after the international filing date. "X" document of particular relevance; the claimed invention			in conflict with the application but principle or theory underlying the	
which i	ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another n or other special reason (as specified)	earmot be considered not involve an inventive step "Y" document of particular re cannot be considered to it	vel or cannot be considered to : o when the document is taken alone elevance; the claimed invention involve an inventive step when the	
other n "P" docume	ent referring to an oral disclosure, use, exhibition or neans ant published prior to the international filing date but can the priority date claimed	document is combined w	with one or more other such docu- a being obvious to a person skilled	
Date of the	actual completion of the international search	Date of mailing of the int		
26	5 July 1994	18 -	08- 1994	
Name and m	nailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk	Authorized officer		
•	Tcl. (+ 31-70) 340-2040, Tx. 31 651 epo nl. Fax: (+ 31-70) 340-3016	Deffner, C-	- A	

INTERNATIONAL SEARCH REPORT

information on patent family members

Inte onal Application No PCT/US 94/02964

Patent document cited in search report	Publication date	Patent mem	family ber(s)	Publication date
EP-A-0293881	07-12-88	US-A- AU-B- AU-A- CA-A- DE-A- JP-A- US-A- US-A-	5187157 623592 1733288 1328332 3878991 1063583 5242904 5250720	16-02-93 21-05-92 08-12-88 05-04-94 15-04-93 09-03-89 07-09-93 05-10-93
US-A-5196948	23-03-93	JP-A- JP-A-	3041478 3041484	21-02-91 21-02-91

Form PCT/ISA/210 (putent family annex) (July 1992)